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(71) Applicant (for all designated States except US): **CAN-  
TION A/S (DK/DK)**; Ørsted's Plads, Building 347,  
DK-2800 Kgs. Lyngby (DK).

(72) Inventor; and

(75) Inventor/Applicant (for US only): **FALTUM, Carsten**  
[DK/DK]; Birkevej 1, DK-3489 Fredensborg (DK).

(74) Agent: **NKT Research & Innovation A/S SCION.DTU**;  
Diplomvej. Bldg. 373, DK-2800 Kgs. Lyngby (DK).

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(54) Title: **SENSOR SYSTEM WITH A REFERENCE SURFACE MIMICKING THE DETECTION SURFACE BUT WITH LOW LIGAND BINDING CAPACITY**

(57) Abstract: The invention concerns a sensor system with at least two flexible units, a sensor unit and a reference unit. The sensor unit comprises a capture surface area functionalised by linking one or more functional groups comprising a capture ligand, such as a member of a specific binding pair. The reference unit comprises an imitated capture surface area which area has been functionalised by linking one or more functional groups, wherein said one or more functional groups linked to the imitated capture surface area of said reference unit do not include a ligand which is identical with said capture ligand. The capture ligand may e.g. be a specific binding partner for a biocomponent, preferably selected from the group consisting of RNA oligos, DNA oligos, PNA oligos, protein, peptides, hormones, blood components, antigen and antibodies. The sensor unit and the method make it possible to reduce the noise, because the signal obtained from the reference unit which is measuring the noise may be subtracted from a signal obtained from the sensor unit.

SENSOR SYSTEM WITH A REFERENCE SURFACE MIMICKING THE DETECTION SURFACE BUT WITH  
LOW LIGAND BINDING CAPACITY

The present invention relates to a sensor system for  
detecting the presence or the amount of a substance e.g.  
5 a target biocomponent in a fluid such as a liquid.

*Bagground of the invention*

It is known from e.g. WO 0066266 and WO 9938007 that  
10 micro-cantilevers can be used for detection of molecular  
interaction. Capture molecules are immobilised on the  
surface of the cantilever. When the capture molecules  
bind to an analyte in the sample that is presented to the  
cantilever, this will induce a change in the surface  
15 stress of the cantilever, and consequently the cantilever  
will deflect and/or stretch.

Measuring the reflection angle from a laser beam that is  
directed to the cantilever can detect a deflection.  
20 Another sensor principle is the use of a piezoresistor  
integrated into the cantilever. In this detection  
principle the deflection/stretch is detected as a change  
in the electrical resistance of the piezoresistor.

25 The signal from the measuring cantilever comprises both  
the signal from the deflection and stretch of the  
cantilever but also from noise.

It has previously been demonstrated that a blank  
30 cantilever can be coupled e.g. in a Wheatstones bridge  
with the measuring cantilever in order to eliminate the  
mechanical noise in the system which may include, but is  
not limited to, external vibration, temperature changes  
etc.

In this case the signal from the reference cantilever can be described as:

$$[2] \quad S_{\text{ref}} = S_{\text{mech. noise}}$$

5

where:

Sref = Total signal from the reference cantilever

The noise of the signal has thereby been reduced significantly. However, the signal measured still includes noise, and it is the object of the present invention to provide a sensor system where the amount of noise is even further reduced than in the prior art solutions described above.

15

#### *Summary of the invention*

It has been found that the signal from the measuring cantilever will be a combination of several components:

20

$$[1] \quad S_{\text{mc}} = S_{\text{spec. bind.}} + S_{\text{mech. noise}} + S_{\text{unspec. bind.}}$$

where:

S<sub>mc</sub> = Total signal from the measuring cantilever

25 S<sub>spec. bind.</sub> = Signal due to the specific binding of the analyte to capture molecules

S<sub>mech. noise</sub> = Signal due to mechanical noise, e.g. external vibrations, temperature variation

30 S<sub>unspec. bind.</sub> = Signal from unspecific binding of molecules in the sample to the capture molecules

Unspecific binding is a substantial problem in all binding assays as it is impossible today to distinguish between specific and unspecific binding. As can be seen 35 the formulas above and the use of a traditional reference cantilever do not eliminate this problem.

The objective of the invention is therefore to provide a sensor system which system does not have the drawbacks as described above. The invention as it is defined in the  
5 claims provides a sensor system with a reduced level of noise compared to prior art sensors.

The sensor system according to the invention comprises at least two flexible units, wherein one of said units is  
10 ``a sensor unit'' and another one is ``a reference unit''. The sensor unit comprises a capture surface area which area has been functionalised by linking one or more functional groups comprising a capture ligand to said capture surface area.

15 The reference unit comprises an imitated capture surface area, which area has been functionalised by linking, preferably covalently linking one or more functional groups. The reference unit differs from the sensor unit  
20 in the composition of the functional groups linked to its surface. As it will be clear from the following, the main issue of the invention is that the reference unit is functionalised but that it contains less or no functional groups which are identical with the capture ligand of the  
25 sensor unit. In one embodiment, the imitated surface area contains no or less members of the specific binding pair which is complementary to the same binding partner member as the capture ligand of the sensor unit.

**Disclosure of the invention**

The sensor system of the invention concerns a set-up as defined in the claims which system comprises a measuring  
5 sensor unit which is coupled to an "intelligent" reference unit. In this set-up it has surprisingly turned out that the reference unit can be used to partly or totally eliminate not only mechanical noise, but also to reduce or eliminate the noise originating from unspecific  
10 binding.

The sensor unit is functionalised by immobilising capture molecules as normally to a capture surface area of the sensor unit. Contrary to the normal procedure, the  
15 reference unit is also functionalised at a surface area designated "an imitated capture surface area".

In one embodiment the noise is reduced significantly, e.g. to 50 % or lower compared to prior art technology.  
20

In one embodiment, the noise resulting from unspecific binding is essentially eliminated i.e. within measuring uncertainty.

25 In the following the term "a ligand" means a type of ligand, and similarly "a binding partner" means a type of binding partner and so on.

Both the capture surface area (which means the capture  
30 surface area of the sensor unit), and the imitated capture surface area (which means the imitated capture surface area of the reference unit) are functionalised by linking one or more functional groups to the surfaces, preferably so that the amount of proteins, amino acids  
35 and/or lipids that binds via unspecific binding to the imitated capture surface area is closer to the amount of

similar components that binds via unspecific binding to the capture surface area than if the reference unit was not functionalised.

- 5 The functional groups linked to the capture surface area and the reference surface area, respectively, may be linked chemically e.g. covalently or ionic bondings; or physically. In one embodiment, one or more of the functional groups are linked via covalent bonds. In one  
10 embodiment, one or more of the functional groups are linked by adsorption. Adsorption means a non-specific physical interaction between the functional groups and the surface area. Adsorption is relatively cheap, easily carried out, and tends to be less disruptive to enzymic  
15 proteins than chemical means of attachment, the adsorption binding being mainly by hydrogen bonds, multiple salt linkages, and Van der Waal's forces. Adsorption bears the greatest similarity to the situation found in biological membranes in vivo and may therefore  
20 in some embodiments be preferred.

The size of the sensor unit capture surface area and the size of the reference unit imitated surface area may differ from each other or it may be equal. If the size of  
25 the sensor unit capture surface area and the size of the reference unit are equal to each other, the calculation of the signal with reduced or no noise is easier than if the sizes differ from each other. In the latter case, a correlation factor compensating for the size difference  
30 should be implemented.

The capture ligand is a member of a specific binding pair. Such ligands which is members of a specific binding pair is well known in the art, and further, information  
35 concerning such binding pair can be found in WO 0066266,

WO 9938007, US 5,156,810, WO 0036419 and WO 9631557 which publication are hereby incorporated by reference.

By the term specific bonding pair is meant any pair of  
5 target molecule/capture ligand with an ability to  
specifically bind to one another e.g. receptor/target  
ligand, enzyme/substrate (or analogue), nucleic acid  
binding protein/nucleic acid etc. Such specific bonding  
pair of target molecule/capture ligand is thereby said to  
10 be complementary to each other.

In one embodiment, the binding pair in the form of the  
capture ligand binding partner and a target binding  
partner is selected among antigen-antibodies or fragments  
15 thereof and nucleic acid strands - nucleic acid strands.

In one embodiment, a molecule that shows similarity to  
the capture molecules is immobilised to the reference  
unit. The molecules on the reference unit though, in this  
20 embodiment do not exhibit a specific binding to the  
analyte that the assay is designed to detect.

In one embodiment, the one or more functional groups  
linked to the imitated capture surface area of the  
reference unit do not include a ligand, which is  
25 identical with the capture ligand. In one embodiment, the  
one or more functional groups linked to the surface area  
of said reference unit do not include a ligand which is a  
member of the specific binding pair.

30

The molecules on the reference unit can belong to any  
class of molecules and do not necessarily have to be of  
the same class as the molecules on the sensor unit.

35 In one embodiment, the functional group linked to the  
imitated surface area of the reference unit includes a

reference ligand. The reference ligand may in one embodiment be present in a number which is 50 % or more such as 75 % or more, such as 90 % or more of the number of capture ligands on the capture surface of the sensor unit. In one embodiment, the reference ligand is of the same chemical class as the capture ligand of the sensor unit. The chemical class may e.g. be one of a) nucleic acids and strands thereof such as DNA oligos, PNA oligos, RNA oligos; b) proteins including peptides, antigen, antibodies and hormones; and c) lipids.

In one embodiment of the sensor system according to the invention, the sensor system is directed to detecting the presence of a preselected target biocomponent. The sensor unit comprises a capture surface area which area has been functionalised by linking a capture ligand, where said capture ligand is a capture ligand for the preselected target biocomponent. The reference unit comprises an imitated capture surface area which area has been functionalised by linking one or more functional groups, wherein the one or more functional groups of the imitated capture surface have less tendency to bind to the preselected target biocomponent than the capture ligand. In one embodiment, the functional groups linked to said imitated capture surface area of said reference unit do not include a ligand, which is a capture ligand for said preselected target biocomponent.

In one embodiment, the amount of proteins, amino acids and/or lipids that binds via unspecific binding to the imitated capture surface area is 50 % or more, such as 60 % or more, such as 70 % or more, such as 80 % or more, such as 90 % or more, such as essentially the same as the amount of similar components that binds via unspecific binding to the capture surface area. In one embodiment, this is measured by using DAKO Human Serum Protein



Calibrator as test medium. In one embodiment, this is measured by using DAKO Lipoprotein (a) Calibrator as test medium.

- 5 When a sample is presented to the sensor unit and reference unit, the analyte, which is a binding partner to the capture ligand (if present), will bind to the capture ligand on the sensor unit. Also various other molecules in the sample will unspecifically bind to the  
10 functional groups on the sensor unit.

In one embodiment, it is desired that the one or more functional groups of the imitated capture surface have a capture tendency of 50 % or less, such as 40 % or less,  
15 such as 30 % or less, such as 20 % or less, such as 10 % or less than the capture tendency of the capture ligand toward said preselected target biocomponent.

In one embodiment, the reference ligand of the reference  
20 unit has essentially the same charge as the capture ligand of at least one sensor unit connected to the reference unit. Essentially the same charge means that the charge of the respective capture/reference ligands differs from each other by 10 % or less, preferably by 5  
25 % or less measured in water.

In one embodiment, the imitated capture surface area of the reference unit has essentially the same pH value as the capture surface area of at least one sensor unit  
30 connected to the reference unit. Essentially the same pH value means that the pH value of the respective surface areas differs from each other by 0.5 pH or less, preferably by 0.1 pH % or less measured in water.

In one embodiment, the reference ligand of the reference unit has essentially the same hydrophility as the capture ligand of the sensor unit.

5 In one embodiment, the reference ligand of the reference unit has essentially the same structure as the capture ligand except for the binding site or sites, which may e.g. be blocked, removed or replaced by non-active chemical group(s).

10

In one embodiment, the imitated capture surface area of the reference unit has a surface tension measured as contact angle to a water droplet which is the same  $\pm 2^\circ$  as the surface tension of the capture surface area of the sensor unit.

15

In one embodiment of the sensor system of the invention the capture ligand is present in a first concentration linked to the capture surface area, and the capture  
20 ligand linked to the imitated capture surface area in a second concentration, wherein said second concentration is substantially less than the first concentration such as 50 % or less, such as 40 % or less, such as 30 % or less, such as 20 % or less, such as 10 % or less than the  
25 first concentration. In this embodiment also a part of the signal originating from a bonding between capture ligand and target ligand will be suppressed, this effect may therefore further be used in determining concentrations of a target biocomponent in a sample.

30

In one embodiment, it is desired that the sensor system according to the invention comprises one or more sensor units and one or more reference units, where at least one, preferably each of the sensor units, is connected to  
35 at least one reference unit. The connection may preferably include a coupling of the connected sensor

unit and reference unit so that a signal obtained from the reference unit is subtracted from a signal obtained from the sensor unit, more preferably said sensor unit and said reference unit being coupled in a Wheatstones  
5 bridge.

The sensor and the reference units are coupled in a way so that the signal from the reference unit (e.g. a cantilever) is subtracted from the signal from the sensor  
10 unit (e.g. a cantilever). This could be done by coupling them in a Wheatstones bridge in the same way as described in WO 0066266 and disclosed in WO 0066266 FIG. 3, the figure and accompanying description hereby being incorporated by reference.

15 The sensor system according to the invention may in principle comprise an unlimited number of sensor units and reference units, and the invention includes embodiments wherein two or more sensor units are coupled  
20 to one reference unit, and embodiments wherein two or more reference units are coupled to one sensor units. The sensor system may preferably include two or more sensor unit, preferably at least 5 sensor units, more preferably at least 10 sensor units, wherein each of said sensor  
25 units preferably comprises a capture surface area which area has been functionalised by linking, preferably by adsorption linking one or more functional groups comprising a capture ligand to said capture surface area, the capture ligand linked to each sensor units being a  
30 member of a specific binding pair. In one embodiment, the capture ligand on one sensor unit preferably is different from the capture ligand of another sensor unit. Thereby several specific target molecules may be detected by the sensor system simultaneously.

35

- In one embodiment, at least one sensor unit and at least one reference unit, which are preferably coupled, have substantially the same size. "Substantially the same size" means within a 10 % variation based on the largest of the sensor/reference units. In one embodiment, "substantially the same size" means within a 5 % variation based on the largest of the sensor/reference units.
- 10 In one embodiment of the system according to the invention, the sensor unit(s) and the reference unit(s), which are coupled together, and preferably all of the sensor units and the reference units may preferably have substantially identical shapes.
- 15 In one embodiment, at least one reference unit coupled to a sensor unit has a thickness which is identical  $\pm 10\%$  of the thickness of the sensor unit.
- 20 In one embodiment, the capture surface area and imitated capture surface area of pair wise sensor/reference flexible units have substantially identical sizes, wherein "substantially identical" should mean within a difference of 20 %, preferably within a difference of 10 %, even more preferably within a difference of about 5 %.
- 25 The flexible unit may in principle have any shape. In one embodiment, the flexible unit or a part of the unit is sufficiently flexible to perform a measurable change due to a stress reaction on the capture surface area/imitated capture surface area when the analytes or substances to be detected are adsorbed or linked to the capture surface area. In one embodiment, the flexible unit should have a free flexing area which is not directly bonded to a solid material.
- 30
- 35

In one embodiment, the flexible units comprise one or more units selected from the group consisting of cantilevers, bridges and membranes. In one embodiment, the flexible units are of micro size, which means that  
5 the flexible units have dimensions which are 500x500x500  $\mu\text{m}$  or less, preferably 5x100x200  $\mu\text{m}$  or less. The shape of the flexible units may e.g. be as described in US 6016686 WO 0066266.

10 In one embodiment, the flexible units comprise a piezoresistor by use of which it is possible to register change in stress of the capture surface area. Further information concerning this aspect can also be found in WO 0066266, which information is hereby incorporated by  
15 reference.

The ligand linked to the sensor unit may preferably be linked via a spacer. Information concerning useful spacer molecules can be found in WO 9631557.

20 In one embodiment, the capture ligand is selected from the group consisting of RNA oligos, DNA oligos, PNA oligos, protein, peptides, hormones, blood components, antigen and antibodies.

25 In one embodiment, the capture ligand is a specific binding partner for a biocomponent, preferably selected from the group consisting of RNA oligos, DNA oligos, PNA oligos, protein, peptides, hormones, blood components,  
30 antigen and antibodies.

The term biocomponent further includes biomolecules and biocomponents selected from the group consisting of tissue, cells, body fluids, blood components,  
35 microorganism, derivatives thereof, or parts thereof.